PATENT SPECIFICATION



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COMPLETE SPECIFICATION

Curing of Polymeric Materials.

We, DAVID AUGUSTINE HARPER and WALTER FAIRBAIRN SMITH, both of Hexagon House, Blackley, Manchester, British Subjects, and Imperial Chemical INDUSTRIES LIMITED, of Chemical House, Millbank, 5 INDUSTRIES London. S.W.1, a Company incorporated under the laws of Great Britain, do hereby declare the nature of this invention and in what 10 manner the same is to be performed, to be particularly described and ascertained in and by the following statement:-

This invention relates to the curing of polymeric materials more especially to 15 the curing of organic discoyanate modified poly-esteramides in admixture with

other polymeric materials.

In Application aumbered 18204/41 (Serial No 580,524) it has been proposed 20 to cure organic discoyanate modified polyester-amides by heating these in the presence of formaldehyde or of a formaldehyde-liberating substance, and pre-ferably, also in the presence of materials 25 which function as curing catalysts, for example; formic, glycollic, oxalic, succinic, maleic, adipic, tartaric, salicylic, anthranilie, phthalie, citric, boric and amearamus, pnraam, curac. borne and phosphoric acids, phthalic and maleic so anhydrides, phthalimide and potassium or sodium dihydrogen phosphate.

In Application numbered 7892/42

(Serial No. 580;526) it has been proposed to cure organic disocyanate modified poly-85 ester-amides by heating these in the presence of formaldehyde or of a formaldehyde-liberating substance and also in the presence of curing catalysts which are substantially neutral but which develop substantially neutral total descriptions, for example butadiene sulphone, 2:3-dimethylbutadiene sulphone, butadiene

methylbutadiene sulphone, tetrabromide, styrenedibromide, acetylene tetrabromide, tribromohydroquinone, 1-45 bromo - 2 naphthol, 1:6 - dibromo-2-

1:4:6 tribremo-2-naphthol, naphthol, methyl-a:β-2:4-dibromo-1-naphthol, β-chloroethyl-α:βdibromopropionate, ethyl a-bromo-prodibromoisobutyrate, 50 pionate, phenyl trichloroacetate. α:α:β trichloropropienitrile, trichloroacetamide, trichloroscetyldiethylamide,

N: Nº-di(trichloroscetyl) acetylanilide,

N: N1-di-(trichloromethylenediamine, acetyl)ethylenediamine, interpolymers of 55 asymmetrical dichloroethylene and vinyl chloride, and chloranil tetrachloro-pbenzoquinone).

In Application numbered 10290/48 (Serial No. 581,146) it has been proposed 60 to use dichromates for curing organic diisocyanate modified polyester-amides.

We have now found that the organic discovanate modified polyester-amides may be cured; with advantage, together 65 with small or large proportions of other polymeric materials which are themselves reactive to formaldehyde and/or di-

chromates. According to the present invention in 70 the curing of organic diisocyanate modified polyester-amides in the manner hereinafter set forth we provide the improvement which comprises curing said modified polyester-amides in uniform 75 admixture with a proportion of a deriva-

tive of cellulose. The invention also comprises heat-curable compositions comprising as the essential ingredients an organic diisocyanate modified polyester-amide, a proportion of a derivative of cellulose, and one or more materials of the kind hereinafter set forth such as are customarily used for curing organic diisocyanate 85 modified polyester-amides.

Water-soluble or water-insoluble derivatives of cellulose include cellulose esters, for example, cellulose nitrate and celluloss acetate, and celluloss ethers such as 90 benzyl cellulose. These materials are modified by the action of formaldehyde thereon, and they are compatible with the organic diisocyanate modified polyesteramides, in the sense that they are capable 95 of forming homogeneous blends there-

The derivative of cellulose may be brought into uniform admixture with the organic diisocyanate modified polyester- 100 amide in several ways. For instance, a derivative of cellulose dissolved in water is slowly added to an organic diisocyanate modified polyester-amide running on a warm rubber mill. Or, a derivative of 105 cellulose in powder form, is milled into an

organic diisocyanate modified polyesteramide on a rubber mill, a sufficiency of water usually being added to render the mix soft. When water is used before or during the incorporation, the mix is usually milled until it is substantially dry, the rolls, if hot, being preferably allowed to cool during the drying so as to minimise the risk of the mix sticking thereto. Alternatively, the components are mixed or milled together (in the absence of water) as such or in the

presence of organic liquids which are solvents for one or more of the materials; the 15 organic liquids are removed as and when convenient. The proportions of the components are

not critical, but usually from about 25 to 175 parts of the derivative of cellulose per 20 100 parts of organic diisocyanate modified polyester-amide are used.

The mixtures are cured by means any of the materials customarily used for curing organic diisocyanate modified polyester amides, that is to say, by heating with a known curing agent, namely, formaldehyde or a formaldehyde liberating substance or a dichromate; preferably in the presence of a known curing cata-30. Lyst, namely, an acid or a material which is substantially neutral but which develops acidity under ouring conditions. These ingredients required for curing are incorporated with the mix whenever con-85 venient but, if water has been used to assist in the formulation of the mix, preferably after the mix is dried.

As well as those already mentioned, one or more additional compounding ingredients may also be used. These include fillers, for example, carbon black, iron oxide, clay, asbestos, blanc fixe, whiting, lithopone and mica; resins, for example, urea-formaldehyde and phenol-formalde-45 hyde resins; other plastic materials, for example, natural or synthetic rubbers,

vulcanised vegetable oils. dark substitute, white substitute, a Cumar resin, wood rosin and pitch; de-tackifying agents, 50 that is to say, materials which reduce the tendency of the mix to stick to the rolls, tentency or see mrx to seate to the rous, for example, steario acid, paraffin wax. coleic, solid, laurio acid and dibutyl ammonium closate; plasticisers, for phithalate, butylphthalyl, butyl glycol-

late, and N-alkyl-toluenesulphonemides; stabilisers (r anti-oxidants, for example, N: N1-hexamethylene-bisortho-hydroxy-benzamide.

N-phenyl-anaphthylamine, N-phenyl-β-naphthyla: a-bis(2-hydroxy-3:5and. dimethylphenyl)butane, as well as others

commonly used in rubber technology. Small quantities of pigments, for example

from 1-3% by weight, such as are customarily used in rubber technology or in the coating composition art may also be used to impart colour. The use of alkaline reacting compounding ingredients should be avoided since these may cause degradation of the polymeric materials.

When the ingredients are mixed, the mix is removed from the mill or mixer, if desired, formed into shapes or spread or 75 calendered on to a substrate, for example . on to the surface of a fabric, or on to the surface of a coated fabric and then curing is effected by heating, for example, in a mould which is preferably in a hydraulic 80 press, or in hot air. Periods of heating varying from a few minutes to several hours at 100-150° C. are usual. If desired, to facilitate shaping or spreading, organic solvents or swelling agents or additional solvents or swelling agents may be incorporated with the materials;

these are removed as and when convenient, Suitable organic solvents acetone, mixtures of benzene and acetone, mixtures of benzene and ethanol, mixtures of benzene and chloroform, mixtures of benzene and methylethyl ketone, mixtures of methyl ethyl ketone and trichlor-

ethylene, and mixtures of acetone and the monosthyl ether of ethylene glycol. Polyesterand

polyamide-forming reactants suitable for making the disccyanate modified polyester-amides to be used for the purposes of the present inven- 100 tion include glycols, for example, ethylene glycol, diethylene glycol, trimethylene glycol, pentamethylene glycol, hearmentylene glycol, dodecamentylene glycol, l. 12-octadecamediol and pents. 105 glycol; aliphatic or aromatic amino-alcohols having at least one hydrogen atom attached to the amino nitrogen atom and preferably containing an aliphatic chain of at least two carbon atoms separat- 110 ing the amino and hydroxyl groups, for propanol; dibasic carboxylic acids or esterforming derivatives thereof preferably adiphatic dicarboxylic acids, for example, 415 malonic, succinic, dutaric, adipic, 9, methyladipic, pimelic, suberic, azelaic undecanedioic, brassylie, isophthalie. hexabydroterephthalic. phenylenediacetic, and acetone-dicarb- 120 oxylic acids; primary and secondary diamines, for example, ethylene diamine, hexamethylenediamine, 8 methylhexamethylene diamine, decamethylenediamine. m-phenylenediamine. N: N1-di- 125 methylhexamethylenediamine.

ethylhexamethylenediamine, and N:N1 dimethyldecamethylenediamine; hydroxymonocarboxylic acids or their ester-forming derivatives, for example, 180

N: N1 di -

6-hydroxycaproic, 10-hydroxyand 12-hydroxystearic acids; decanoic polymerizable monoaminomonocarboxylic acids, or their ester-forming derivatives, 5 for example, 6-aminocaproic acid or its caprolactam, and 9-aminolactam. nonanoic, 11-aminoundecanoic and 12-

aminostearic acids. The polyester-amides are made in

10 known manner by heating the selected reactants at polymerizing temperatures, usually in the absence of air or oxygen, under conditions whereby water is removed from the reaction mixture. When 15 a diamine is to be used, it is conveniently used in the form of the corresponding di-

ammonium salt from some of the dibasic carboxylic acid to be used.

The polyester-amides are modified with 20 organic diisocyanates in known manner for example, by mixing them, for example by stirring, milling or kneading, with the organic diisocyanate and then heating the mixture, for example, to a 25 temperature of 100-200° C. for a period of 10-720 minutes. Up to about 10 percent; usually 8-7%, by weight of the diisocyanate is used.

Examples of organic discovanates so includes ethylene discovanate, tritrimethylene diisocyanate, tetramethylene diisocyanate, hexamethylene diisocyanate decamethylene diisocyanate, p-phenylene diisocyanate, m-phenylene diisocyanate, 85 p:p¹-diphenyl diisocyanate, diphenylmethane-4:41-diisocyanate, naphthalene

diisocyanates and adipyl diisocyanate. · The new heat-curable compositions of the invention may be used in the fabrica-40 tion of a variety of articles, in which they may or may not be supported on a substrate and/or interspersed with fillers.

For instance, they may be used in the construction of organic liquid resistant 45 articles of all kinds, for example, gaskets, packings, hose, diaphragms for pumps and the like, as well as in the fabrication of flexible containers. They may also be used in the coating of rollers, blankets and 50 stereos for use in the printing industry, or to provide protective sheathings for insulated electric cables and other elec-

trical conductors. They also find appliestion in the coating of the balls for games, 55 tyres and flexible materials generally, including fabrics, protective clothing, leather cloth and floor coverings, and generally in the construction of articles

requiring the use of a material having 60 physical properties resembling those of rubber, but also having a good resistance to the action of organic fluids and a low permeability to gases and vapours.

They are well adapted for application 65 in the form of lacquers or finishing com-

positions for all kinds of surfaces. They may be formed into films or sheets, for example, as substitutes for leather or as wrapping films, and they may be used as adhesives for a wide variety of materials, for example, wood, metals, fabrics, paper, leather and regenerated cellulose

The invention is illustrated but not limited by the following Examples, in which the parts and percentages are expressed by weight, unless otherwise stated:---

EXAMPLE 1.

Into 100 parts of an organic diisocyanate modified polyester-amide there are milled 10 parts of titanium dioxide, 5.0 parts of hexamethylolmelamine hexamethyl ether and 0.75 parts of 2:4-dichloro I-naphthol. The mix is then let down in 100 parts of a mixture of equal volumes of benzene and acetone, and 125 parts of a 20% solution of cellulose acetate (acetone-soluble) are stirred in. The mix is thinned with acetone to a solids content of 10-15%, and

is ready for use. The composition is flowed on to a glass plate, the solvent is allowed to evaporate, and the so obtained film is cured by heat-

ing in air at 125° C. for 2 hours.

The cured film is tough and has a rubbery handle; its surface is very

resistant to scratching.

When the recipe of the Example is repeated using twice the quantity of the solution of cellulose acetate, a film is 100 obtained which is tougher, less rubbery and extremely resistant harder. scratching.

Similarly, when the recipe of the Example is repeated using three times the 105 quantity of the solution of cellulose acetate, there is an increase of toughness and hardness, and a decrease of rubbery properties; the film has outstanding resistance to scratching.

EXAMPLE 2. Into 100 parts of an organic diisocyanate modified polyester-amide there are milled 10 parts of titanium dioxide, 5.0 parts of hexamethylolmelamine hexa- 115

methyl ether and 0.75 parts of 2:4-dichloro-1-naphthol. The mix is let down in 100 parts of a mixture of equal volumes of benzene and acetone, and 250 parts of a 20% solution of nitrocellulose 120 obtainable commercially under the name

"Collodion Cotton HX 30/50" are stirred in. The mix is thinned with acetione to a solids content of 10-15%. and is ready for use

The composition is flowed on to a glass plate, the solvent is removed by evaporation, and the so obtained film is cured by heating in air at 125° C. for 2 hours.

The cured film is tough, rubbery and 130

has a very good scratchi resistance.

When the recipe of the Example is repeated using half as much again of the nitrogellulose solution, a tougher, less rubbery film with better scratch resistance is obtained.

Example: 8.

A lacquer is prepared by mixing together 100 parts of an organic 10 discovyanate modified polyester-awaide, 60 parts of: the cellulose acetate used im Example 1, 10 parts of carbon black, 5 parts of hexamethylol-melamine hexamethyl ether, 0.75 parts of ethylene glycol monoethyl ether and 350 parts of eccent

The so obtained lacquer is used to provide a top-coating for the hydrolysed.
20 leather-organic discognants modified polyaster-amide coated, fabric obtained in the manner set forth in Application num-

bared 21389/43 (Serial No. 583,862), and more specifically as follows:—

180 parts of disintegrated scrap vegetable-tanned leather are mixed with sufficient cold water to form an easily stirred sturry, the slurry is heated up to boiling and boiled for about 5 minutes. The leather settles in the form of a crumbly mass from which the liquid is poured off. The wet mass is added in small amounts to 100 parts of an organic diisocyanate modified polyester-amide running 35 on to a rubber mill with the rolls heated to about 709 C.; the mix is milled until homogeneous, and then dried on the rolls. The following ingredients are then added in the order listed, 0.5 parts steam acid, 40 30 parts of titanium dioxide, 5 parts of hexamethylolmelamine hexamethyl ether and 0.75 parts 2:4-dichloro-1-naphthol, milling is continued until the mix is again homogeneous, and it is then sheeted off

45 the mill.

The sheet is added to its own weight of a benzene-ethanol mixture (75:25 by volume) in a Werner Pfleiderer mixer, and mixed until a smooth dough is

obtained. The dough is spread on to a 50 cotton twill fabric, the organic liquids are allowed to evaporate.

The coated spreading is then cured by heating in air at 125—130° C. for 2 hours.

The cured material has a high gloss, an excellent abrasion, somb and flex resistance, and an outstanding resistance to scratching or marking.

The organic diisocyanate modified polyester-amide used in the above of Examples is that described in Example 7 of Application numbered 18204/41 (Serial No. 580,524).

Having now particularly described and ascertained the nature of our said invention and in what manner the same is to be performed, we declare that what we claim

is:—

1. In the process of curing organic diisocyanate modified polyester-amides in the known manner hereinbefore set forth, the improvement which comprises curing said modified polyester-amides in uniform mixture with a proportion of a derivative

of cellulose.

2. The improvement which comprises curing organic diisooyanate modified polyesta-amidés in uniform admixture with a proportion of a derivative of cellulose in the manner hereinbefore pasticu-

lanly described and assertained especially with reference to the foreging Examples.

3: Organic discovanate modified polyester-amides whenever cured in unform admixture with a proportion of a federivative of cellulose according to either

of the preceding claims.

4. Hest-curable compositions comprising as the essential ingredients an organic
discoyanate modified polyester-amide, a
preportion of a derivative of cellulose, and
one or more materials of the kind hereinbefore set forth such as are customarily
used for ouring organic discoyanate

modified polyester amides.

Dated the 26th day of September, 1945.

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Solicitor for the Applicants.